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(73) Proprietor: **KIMBERLY-CLARK CORPORATION**
401 North Lake Street
Neenah,
Wisconsin 54956-0349 (US)(72) Inventor: **Bell, Anita Salina**
616 Belcourt Parkway
Roswell, GA, 30076 (US)
Inventor: **Cohen, Bernard**
381 Lakeshore Drive
Berkley Lake, GA 30136 (US)
Inventor: **Morman, Michael Tod**
555 Kings Peak
Alpharetta, GA, 30202 (US)(74) Representative: **Diehl, Hermann O. Th., Dr. et**
al
Diehl & Glaeser, Hiltl & Partner
Patentanwälte
Postfach 19 03 65
D-80603 München (DE)

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Description

The present invention relates to a method of preparing a high hydrohead fibrous porous web material that increases the retentive water absorbency of the web. The retentive water acquisition rate may also be increased. As used herein the term "absorbency" generally refers to the ability of a material to acquire a fluid and the acquisition rate refers to the rate of such acquisition. An example of a use where high absorbency and high acquisition are desired would be wiper type materials. In addition to having the characteristics of high absorbency and high acquisition rate, wipers desirably also should have the characteristic of high retentive absorbency and high retentive acquisition rate. The term retentive acquisition rate is used herein to designate comparison of the rate of acquisition of a fluid by a material when the material is first used to acquire the fluid as compared to the second, third and fourth times the the material is used to acquire the fluid. Improved retentive acquisition rate is evidenced by smaller decreases in the rate of acquisition with multiple uses. Likewise, the term retentive absorbency is used to designate comparison of the amount of fluid acquired by a material when the material is first used to acquire the fluid as compared to the amount of fluid acquired when the material is used a second, third, fourth time to acquire the fluid. Improved retentive absorbency is evidenced by smaller decreases in the amount of fluid absorbed by the material with multiple uses. In other words, the ability of the material to reabsorb fluid after having, in our test, been exposed to fluid, wrung out and allowed to dry.

The inventors found that the application of corona discharge treatment to low hydrohead webs whose surface includes a surface active agent having a hydrophile-lipophile balance of 6 or greater results in a significant increase in the retentive wettability, as defined therein, of such webs. Low hydrohead webs of that type would generally be unsatisfactory for use as a wiper material due to their open pore structure which would greatly reduce the ability of the web to acquire fluids. Conversely, high hydrohead webs, as defined herein, would generally be undesirable for use in applications where rapid transmission of large amounts of fluid through the material is desired. This undesirability arises from the generally tight, closed pore structure of high hydrohead materials. Such pore structure would inhibit the passage of fluids therethrough in rapid fashion.

In the past, hydrophobic wipers have been subjected to treatment with surfactants to improve their characteristics. The wipers have been treated with surfactant by (1) passing the formed wiper through a bath containing the surfactant in either neat or solution form and drying the wiper as needed so that a given amount of the surfactant is deposited on the wiper, or (2) spraying a surfactant in either neat or solution form on the fibers as they are being formed or on the fibrous porous web and drying the wiper as needed so that a given amount of the surfactant is deposited on the wiper, or, (3) adding surfactant to a thermoplastic resin prior to extrusion and formation of the resin into a thermoplastic porous web material. In the later situation, under known process conditions, the added surfactant exudes or migrates to the surface of the fibers of the porous web material during or shortly after fiber formation. This phenomenon has been referred to as "blooming" the surfactant. It is believed that blooming results from the insolubility of the surfactant in the thermoplastic polymer as the polymer cools. See US-A-4,535,020 to Thomas et al - (hereafter Thomas et al 020) which demonstrates surfactant blooming in a diaper liner formed from a perforated film.

A wiper made from a hydrophobic material, such as a thermoplastic polymer, will not readily acquire or absorb spilled fluids because the surface tension of the fluid is greater than the critical surface energy of the hydrophobic material. Surface tension is the contractile surface force of a fluid where the fluid tries to assume a spherical form and to present the least possible surface area. It is usually measured in dynes per centimeter. Accordingly, because of its effect on the insulating fluids, surfactant has been previously applied to wipers. Application of a surfactant onto a wiper material may make a nonabsorbing wiper absorbant by at least two mechanisms: (1) Surfactants present on the wiper can dissolve into a fluid and lower the surface tension of the resulting solution to more equal the critical surface energy of the wiper material. Accordingly, when a surfactant coated wiper is used to wipe up a fluid such as water, the surfactant acts to lower the surface tension of the fluid and allow the fluid to be acquired at a faster rate and for a larger amount of fluid to be absorbed into the wiper. In this situation, a certain amount of the surfactant on the wiper is lost with each wiping and wringing and unacceptable acquisition rate and absorbency occurs at some following wiping due to the lack of availability of surfactant to lower the surface tension of the fluid. (2) The surfactant can be coated onto the fibers making up the wiper, making the fiber surface of the wiper more hydrophilic, i.e., increase the apparent critical surface energy of the fibers. In this situation the wiper would have permanent absorbency if the surfactant did not dissolve in the fluid the wiper was used to pick up.

As any anyone will testify, it is an aggravating event when a disposable wiper fails in its appointed task of rapidly acquiring and absorbing a fluid spill.

Accordingly, it has been a goal of those in the art to provide a high hydrohead porous web wiper material which has an improved acquisition rate and absorbency. This was the initial goal because, if the material cannot acquire and absorb fluid at all, the material cannot function as a wiper. Additionally, it has been a goal of those in the art to provide an high hydrohead porous web wiper material which has an improved retentive acquisition rate and improved retentive absorbency. That is, when dried and wrung-out between wipings, the wiper has a significant increase in the number of times it can be used to absorb fluid. This goal is desirable not only from the standpoint of allowing a given disposable wiper to be used more times but also from an environmental standpoint in that fewer wipers will be disposed into the environment.

Corona discharge treatment of films is old in the art and it is known that corona discharge treatment of a polymer film in the presence of air entails substantial morphological and chemical modifications in the polymer film's surface region. See Catoire et al, "Physico-chemical modifications of superficial regions of low-density polyethylene (LDPE) film under corona discharge," Polymer, vol. 25, p. 766, et. seq, June, 1984.

Generally speaking, corona treatment has been utilized to either (1) improve the print fastness on the film, or (2) to perforate the film. For example, US-A-4,283,291 to Lowther describes an apparatus for providing a corona discharge, and US-A-3,880,966 to Zimmerman et al discloses a method of using a corona discharge to perforate a crystalline elastic polymer film and thus increase its permeability. US-A-3,471,597 to Schirmer also discloses a method for perforating a film by corona discharge. US-A-3,754,117 to Walter discloses an apparatus and method for corona discharge treatment for modifying the surface properties of thin layers or fibers which improve the adhesion of subsequently applied inks or paints or of subsequent bonding.

It also is possible to treat a diaper liner material with a corona discharge and then immediately dip the film in a surfactant solution. Because the corona effect on the material generally starts to immediately decay, it is important to get the corona treated material into the bath as quickly as possible. Such a method is discussed in Japanese KOKAI Patent JP-A-63[1988]-211375. This document discloses a method for producing a nonwoven fabric having a long lasting hydrophilicity. The method involves first treating a nonwoven fabric of synthetic fiber by a corona discharge and then coating the treated fabric with about 2-10 grams per square meter of fabric of surface active agent.

Of particular interest is the fact that Thomas et al 020 is directed to the utilization of corona discharge in conjunction with surfactant treated films to effect improved wettability, i.e. higher fluid transmission rates and therefore decreased run-off of fluid. In this regard Thomas et al 020 states that a perforated film which has been treated with surfactant and which is then corona discharge treated results in a film with very low, zero or near zero fluid run-off on the first run-off test. Thomas et al 020 reports that this effect is accomplished because the corona discharge treatment acts on the chemical additive, the surfactant, to provide the perforated film with a zero or near zero percent run off. Thomas et al 020 postulates that this effect is achieved due to the surfactant providing a greater polarizability to the film than the film would have without the surfactant being added. The corona discharge treatment provides additional polarizing effect and, in combination with the surfactant, provides improved wettability. Because Thomas et al 020 is directed toward use of the perforated film as a diaper liner, it does not appear to address the questions of acquisition rate and absorbency. Acquisition rate, as defined herein, usually does not apply to a film and diaper liners are generally designed to be permeable to fluids as opposed to absorbing them. Lastly, Thomas et al 020 does not appear to address retentive capabilities at all because the testing reported therein is directed to one-time exposure to fluid.

The object of the present invention is to overcome the above mentioned drawbacks of the prior art. Further objects of the present invention will become apparent from the details given below.

These objects are solved by the method of independent claim 1 and the product of independent claim 6. Preferred embodiments of the method of claim 1 are given in dependent claims 2 to 5. Further advantageous features of the methods and products of the invention are obvious from the description and the drawings.

Accordingly, the present invention provides a method whereby the retentive averaged normalized water absorbency of high hydrohead porous webs is improved.

Furthermore, the present invention provides a high hydrohead fibrous porous web material having an increased retentive averaged normalized rate of water acquisition.

In response to the above, we have devised a method of treating a high hydrohead fibrous porous web material to increase the web's retentive water acquisition rate (averaged normalized rate of water absorption in subsequent reabsorptions as compared to the initial absorption rate) and retentive water absorbency (averaged normalized amount of water absorbed in subsequent reabsorptions as compared to the amount initially absorbed). The method generally includes the steps of: (1) providing a high hydrohead fibrous

porous web having a surface concentration of at least about 0.05 percent, by weight of the web, of a surface active agent having a hydrophile-lipophile balance of at least about 6; and (2) applying a corona discharge equivalent to a charge of at least about 8.6 Wmin/m² (about 0.8 watt min./ft²) per side of the web to the surface active agent bearing web. The resultant web will have a percent decrease in the averaged normalized water absorbed, at two minutes, of less than about 50 weight percent in each of the second, third and fourth times the material is tested in accordance with absorbency test A when compared to the averaged normalized water absorbed upon being initially tested in accordance with absorbency test A. In some embodiments, the web will have an average absorbency decrease, as defined above, of less than about 25 percent.

In some embodiments the resultant web will have a percent decrease in the averaged normalized water absorbed, at one minute, of less than about 50 weight percent in each of the second, third and fourth times the material is tested in accordance with absorbency test A when compared to the averaged normalized water absorbed upon being initially tested in accordance with absorbency test A. Additionally, in some embodiments, the resultant web will have a percent decrease in the averaged normalized water absorbed, at one minute, of less than about 25 weight percent in each of the second, third and fourth times the material is tested in accordance with absorbency test A when compared to the averaged normalized water absorbed upon being initially tested in accordance with absorbency test A.

The treated webs generally also have improved retentive averaged normalized rates of water absorption. Thus, generally, the resulting webs have a percent decrease in the averaged normalized rate of water absorbed, in the first 2.4 seconds of absorption, of less than about 50 percent in each of the second, third and fourth times the material is tested in accordance with absorbency test A when compared to the averaged normalized rate of water absorbed upon being initially tested in accordance with absorbency test A. For example, the webs may have such improved retentive averaged normalized rates of water absorption that the averaged normalized rate of water absorbed, in the first 2.4 seconds of absorption, decreases less than about 25 percent in each of the second, third and fourth times the material is tested in accordance with absorbency test A when compared to the averaged normalized rate of water absorbed upon being initially tested in accordance with absorbency test A. Even more particularly, the webs may have such improved retentive averaged normalized rates of water absorption that the averaged normalized rate of water absorbed, in the first 2.4 seconds of absorption, decreases less than about 10 percent in each of the second, third and fourth times the material is tested in accordance with absorbency test A when compared to the averaged normalized rate of water absorbed upon being initially tested in accordance with absorbency test A.

From about 0.05% to about 3%, by weight of the web material, of surface active agent may be adhered to the web material. For example, from about 0.1% to about 1%, by weight of the web material, of surface active agent may be adhered to the web material. More particularly, from about 0.1% to about 0.4%, by weight of the web material, of surface active agent may be adhered to the web material. Even more particularly, from about 0.2% to about 0.3%, by weight of the web material, of surface active agent may be adhered to the web material.

The equivalent of from at least about 8.6 Wmin/m² (about 0.8 watt min./ft²) per side of the web material of corona discharge may be applied to the web material. Preferably the equivalent does not exceed about 161.5 Wmin/m² (about 15 watt min./ft²). More preferably, the equivalent of from about 10.7 to about 107.6 Wmin/m² (about 1 to about 10 watt min./ft²) per side of the web material of corona discharge is applied to the web material. More particularly, the equivalent of from about 21.53 to about 86.11 Wmin/m² (about 2 to about 8 watt min./ft²) per side of the web material of corona discharge is applied to the web material.

In one embodiment the process includes the steps of (1) forming a melt from a thermoplastic fiber forming material; (2) adding, to the melt, an amount of surface active agent having a hydrophile-lipophile balance of at least about 6 sufficient to effect a surface concentration of the surface active agent of at least about 0.05%, by weight of the resulting fibrous porous web material; (3) forming the melt into fibers and the fibers into a high hydrohead fibrous porous web under conditions which allow at least 0.05%, by weight of the fibrous porous web, of the surface active agent to bloom to the surface of the fibers of the porous web; and (4) applying a corona discharge equivalent to a charge of at least about 0.8 watt minute per square foot of the porous web to the surface active agent bearing web material.

Because not all of the surface active agent added to the melt blooms, the amount of surface active agent added to the melt is generally greater than the amount desired to be present on the surface. Accordingly, the amount of surface active agent added to the melt may vary with the surface active agent used, the thermoplastic material used to form the web and/or the process conditions of forming the web.

As is the case generally, in this embodiment the equivalent of from about 8.6 to about 161.5 Wmin/m² (about 0.8 to about 15 watt min./ft²) of the web material of corona discharge may be applied to the web

material. For example, the equivalent of from about 10.7 to about 53.8 Wmin/m² (about 1 to about 5 watt min./ft²) of the web material of corona discharge is applied to the web material. More particularly, the equivalent of from about 21.53 to about 43 Wmin/m² (about 2 to about 4 watt min./ft²) of the web material of corona discharge is applied to the web material.

5 As used herein, the term 'surface active agent' refers to any compound that reduces surface tension when dissolved in water or water solutions or which reduces interfacial tension between two liquids or a liquid and a solid. Specifically according to the present invention there may be used any of the three general categories of surface active agents, namely detergents, wetting agents (i.e. surfactants) and emulsifiers alone or in combination.

10 Especially preferred in all embodiments of the invention are surface active agents selected from the group including one or more wetting agents, emulsions and dispersants.

In all embodiments the hydrophile-lipophile balance of the surface active agent will be about 6 or greater. For example the hydrophile-lipophile balance may range from 6 to about 20. More particularly, the hydrophile-lipophile balance of the surface active agent may range from 8 to about 20. Even more
15 particularly, the hydrophile-lipophile balance of the surface active agent may range from 10 to about 20.

The present invention is also directed to products preparable by the process. That is, the invention is generally directed to a fibrous porous web which has a high hydrohead when tested in accordance with Test A prior to surfactant and corona treatment in accordance with the invention and which has improved retentive averaged normalized absorbency and improved retentive averaged normalized water acquisition
20 rates after surfactant and corona treatment.

The fibrous porous web material may include a polyolefin or a blend of polyolefins or any other suitable material which may be formed into a fibrous porous web. For example, the fibrous porous web may be formed from polyethylene or polypropylene.

The fibrous porous web material may be formed by any of the wide variety of processes which provide
25 a high hydrohead fibrous porous web. For example, the fibrous porous web may be formed by meltblowing so that the fibrous porous web includes meltblown fibers.

Figure 1 is a schematic representation of one process for carrying out the present invention.

Figure 2 is a schematic representation of a second process for carrying out the present invention.

As used herein the term "high hydrophile-lipophile balance" refers to a surface active agent having a
30 hydrophile-lipophile balance of about six (6) or greater.

As used herein the term "surface active agent" refers to any compound that reduces surface tension when dissolved in water or water solutions or which reduces interfacial tension between two liquids, or between a liquid and a solid. There are three general categories of surface active agents: detergents, wetting agents (i.e. surfactants) and emulsifiers.

35 The term "hydrophile-lipophile balance" (HLB) is well known to those in the art. The HLB of a nonionic surfactant is the approximate weight percent of ethylene oxide in the surfactant divided by 5. The numerical scale of HLB values ranges from 1 (completely lipophilic or oil-loving) to 20 (completely hydrophilic, or water-loving). Refer to W.C Griffin, J. Soc. Cosmetic Chemists 317-326 (1949). In some instances the HLB of a material is determined by comparing its activity to known materials having known HLB's.

40 As used herein the term "high hydrohead material" refers to a porous web material which supports more than about 25 centimeters of water when its hydrohead is measured in accordance with Method 5514 - Federal Test Methods Standard No. 191A. In all cases the hydrohead of the porous web material is determined by measurement either before the web has been treated with surface active agent and corona discharge as is required by the present invention or, if such is not possible, after extraction of the surface
45 active agent from the web.

As used herein the term "water absorbency" refers to the amount, in grams, of water that a 7.62 cm by 20.32 cm (3" x 8") sample (folded as described in Test A, below) of high hydrohead porous web material can vertically acquire within a given amount of time.

50 As used herein the term "normalized water absorbency" refers to the calculated amount, in grams, of water per gram of web that a one gram sample of high hydrohead porous web material can vertically acquire within a given amount of time. This value is calculated by multiplying the "water absorbency" value for a given time period by (1/the weight of the sample).

As used herein the term "averaged normalized water absorbency" refers to the average of three "normalized water absorbency" replicates of the material treated in accordance with the invention. In the
55 example, the "averaged normalized water absorbency" value of the non-corona treated material was attained by averaging four replicates.

As used herein the term "rate of water absorbed" (rate) refers to the rate, in grams per second, of vertical water acquisition of a 7.62 cm by 20.32 cm (3" by 8") sample (folded as described in Test A, below)

of high hydrohead porous web material within a given amount of time.

As used herein the term "normalized rate of water absorbency" refers to the calculated rate, in reciprocal seconds, that a one gram sample of high hydrohead porous web material can vertically acquire within a given amount of time. This value is calculated by multiplying the "rate of water absorbed" (rate) value for a given time period by (1/the weight of the sample, in grams).

As used herein the term "averaged normalized rate of water absorbed" refers to the average of three "normalized rate of water absorbed" replicates of the material treated in accordance with the invention. In the example, the "averaged normalized rate of water absorbed" value of the non-corona treated material was attained by averaging four replicates.

All absorbency and rate of acquisition data given herein were obtained through the use of Water Absorbency Test A, hereinafter Test A. The purpose of absorbency Test A is to quantitatively measure the absorbency and rate of acquisition properties of a porous fibrous web such as a nonwoven web.

Test A requires the following materials/equipment: (1) samples of materials to be tested cut in 7.62 cm by 20.32 cm (3" x 8") size; (2) staples; (3) distilled water; (4) one 250 ml. beaker; (5) one small lab jack; (6) an Instron model 1122 with strip recorder; (7) a Lab Wringer, a #LW838 Atlas Electric Devices Co. of Chicago Ill, was used; (8) one 500 gram load cell for the Instron and (9) one standard ten gram weight.

Sample preparation for Test A is as follows: (1) 7.62 cm by 20.32 cm (3" x 8") samples of the material to be tested are obtained; (2) the sample is folded in on itself lengthwise 2.54 cm (one inch) from one side; (3) the sample is folded in on itself lengthwise 2.54 cm (one inch) from the other side to produce a three ply 2.54 cm by 20.32 cm (1" x 8") sample; (4) the sample is folded widthwise in half; and (5) the sample is stapled 0.3175 cm (1/8 inch) from the widthwise fold. The resultant sample is a butterfly configuration with each "wing" having three piles of sample material.

In order to conduct Test A, the Instron must first be prepared. This is done by installing the 500 gram load cell in the Instron and calibrating the machine with the 10 gram weight. The strip recorder should read 0 to 10 grams (2.54 cm (1 inch) per gram). Next the lower jaws are removed from the Instron and replaced with a lab jack. The beaker which is filled with distilled water is placed on the lab jack. The side of the beaker is marked to record the height of the water in the beaker. It is important that this level be maintained at as constant a level as possible.

Placement of a sample in the Instron should be consistent and is accomplished as follows: (1) a start-up sample is placed in the upper jaws of the Instron with the stapled end down; (2) the lab jack is used to raise the beaker so that the level of the water will be one-eighth inch above the staple (the folded edge of the sample will be one-fourth inch below the surface of the water); and (3) the height of the jack is recorded. It is important that the beaker be raised to the same height for each test.

Sample testing is accomplished as follows: (1) a sample to be tested is placed in the jaws of the Instron as stated above; (2) the strip recorder of the Instron is started and allowed to run for ten seconds to obtain a reading of the sample weight; (3) the level of fluid in the beaker is checked to ensure that it is at the mark that has been placed on the beaker; (4) the lab jack is used to raise the sample to the same height as was recorded with the start-up sample [this step should be done quickly and smoothly to minimize irregularities in the climbing portion of the curve]; (5) the test is allowed to proceed for three minutes: a chart speed of 12.7 cm (5 inches) per minute was used in all cases; (6) once the three minutes has elapsed, the recorder is turned off, the lab jack is used to lower the beaker and the sample is removed from the jaws of the Instron; (7) the staple is carefully removed from the sample but the sample is maintained in its six-ply configuration; (8) a lab wringer is used to remove excess water from the sample; [13.6 kg (30 pounds) added to the wringer arm is adequate] (9) after the sample is put through the wringer, it is unfolded and allowed to dry [5 hours is ample for a 67.82 g/m² (2 oz/yd²) meltblown sample].

The data obtained in test A are as follows: (1) total sample weight is the value read from the baseline of the Instron recorder plot. [The scale of the paper in these tests was 1 inch per gram with a zero to ten gram range.]; (2) actual sample weight is the value calculated to be the total sample weight minus the weight of the staple used to hold the sample fold intact; (3) the water absorbed value is read as the gram weight absorbed amount recorded at 1.2 seconds, 2.4 seconds, 1 minute and 2 minutes of elapsed time. Early points are used to calculate acquisition rate; later points are used to compare overall absorption capacity. Total water absorbed is calculated to be the difference between the baseline total sample weight and the weight read from the curve for a given time. [Note: If the weight on the curve is less than the baseline weight, the amount of water absorbed is recorded as zero. This occurs as the result of a buoyant effect as the acquisition rate decreases.] (4) the rate is the value of the slope of the climbing portion of the curve and is calculated by linear regression using water absorbed readings for early points, i.e. "the points (0 sec., 0 grams), (1.2 sec., Y₁ grams) and (2.4 sec., Y₂ grams). Note that Y₁ = water weight at 1.2 seconds (weight absorbed at 1.2 seconds minus total sample weight) and Y₂ = water weight at 2.4 seconds (weight

absorbed at 2.4 seconds minus total sample weight).

All data have also been normalized and given in terms of grams of water absorbed per gram of tested material. Actual sample weight were used in these calculations.

As used herein the term "decrease in averaged normalized rate" refers to the percentage decrease in the rate of water absorption of a given sample in its second, third, and fourth times of testing, in accordance with Test A, as compared to the rate of water absorption calculated in its first time of testing when done in accordance with test A. Any increase in the rate is reported as a zero decrease.

As used herein the term "decrease in averaged normalized water absorbed" refers to the percentage decrease in the amount of water absorbed by a given sample in its second, third, and fourth times of testing, in accordance with Test A, as compared to the amount of water absorption calculated in its first time of testing when done in accordance with test A. For consistency, the point in time of measurement of the amount of water absorbed must be the same. Thus, this data can be reported at, for example, one minute, two minutes or any other convenient time. The values are reported at 1 and 2 minutes herein.

Referring now to the drawings where like reference numerals represent like structure or like process steps and, in particular, to Figure 1 which schematically illustrates apparatus 10 for forming and treating a high hydrohead fibrous porous web material to improve the retentive water absorbency and retentive water acquisition rate of the material. The process may be initiated by supplying pellets (not shown) of a fiber-forming thermoplastic material which may be, for example a polyolefin or a blend of polyolefins such as polypropylene or polyethylene into the hopper 12 of an extruder 14.

While any thermoplastic fiber forming material may be useful, one desirable material is a polypropylene which may be obtained from the Shell Chemical Company under the trade designation 5A09. The shell 5A09 polypropylene has a melt flow rate of about 40 decigrams per minute when measured in accordance with ASTM D 1238 at 230 °C.

Many other thermoplastic polymers are suitable for use as the fiber forming polymer. Specific, non-limiting examples of such polymers include: polyolefins such as low density polyethylene, linear low density polyethylene and high density polyethylene. The materials may be plasticized with suitable plasticizers, and other additives known in the art may be added to achieve the desired physical characteristics.

Elastomeric polymers may be used to form the fibrous porous web. Such polymers include: polyester elastomeric materials, polyurethane elastomeric materials, polyetherester elastomeric materials, polyamide elastomeric materials, and the various elastomeric A-B-A' block copolymer materials disclosed in US-A-4,663,220 to Wisneski et al, which is hereby incorporated by reference.

Neat or a solution of a surface active agent is sprayed onto the fibers as they are formed or on the formed web 22 from a spraying apparatus which may be a spray boom 19. The surface active agent may be, for example, an emulsion, a wetting agent or a detergent having a hydrophile-lipophile balance of at least about 6 or greater. The surface active agent may be nonionic, cationic or anionic. If the surface active agent is nonionic, it is desirable that it have at least 3 ethylene oxide groups. One desirable surface active agent as a surfactant is Na-di(2-ethylhexyl) sulphosuccinate, which may be obtained from American Cyanamid under the trade designation Aerosol OT. Aerosol OT has an equivalent hydrophile-lipophile balance of greater than about 13. It has been reported that the hydrophile-lipophile balance of Aerosol OT is about 13.5. See, US-A-4,013,863 to van Osenbruggen, et. al. at Table I, therein, and US-A-3,904,728 to Davis, et. al. Another surface active agent which may be used may be obtained from the Rohm & Haas Company under the trade designation Triton X-102. Rohm & Haas literature states that the X-102 is a nonionic octylphenol liquid surfactant having from 12-13 ethylene oxide units. The material is about 73%, by weight, ethylene oxide, has a Brookfield viscosity at 25 °C. (12 rpm) of 330, and has a calculated hydrophile-lipophile balance of about 14.6. Other Triton brand name materials may be utilized in the present invention. Exemplary of which are Triton X-35 which is a nonionic octylphenol series material having three ethylene oxide units and a calculated hydrophile-lipophile balance of 7.8; Triton RW 50 which is a cationic material, $(t-C_{12-14}NH(CH_2CH_2O)_5H)$, having an average of five ethylene oxide units and a measured hydrophile-lipophile balance of 12-14; Triton RW 100 which is a cationic material, $(t-C_{12-14}NH(CH_2CH_2O)_{10}H)$, having an average of 10 ethylene oxide units and a measured hydrophile-lipophile balance of 16; Triton DF 12 which is a nonionic modified polyethoxylated alcohol that has a calculated hydrophile-lipophile balance of 10.6 and Triton DF 18 which is a nonionic biodegradable modified alcohol that has a calculated hydrophile-lipophile balance of 11.3.

It is desirable for the surface concentration of the surface active agent on the surface of the fibers of the web to be at least about 0.05 weight percent of the web. For example, from about 0.05 percent, by weight, to about 3 percent, by weight of the web. More particularly, from about 0.10 percent, by weight, to about 1.0 percent, by weight of the web. For example, from about 0.1 percent, by weight, to about 0.4 percent, by weight, of the web. Even more particularly, from about 0.20 percent, by weight, to about 0.30 percent by

weight of the web. In one embodiment the surface concentration is about 0.30 percent by weight of the web 22.

Because not all of the sprayed surface active agent remains on the fibers, the amount of surface active agent applied to the fibers is generally greater than the amount desired to be present on the surface. Accordingly, the amount of surface active agent sprayed on the fibers may vary with the surface active agent used, the thermoplastic material used to form the web and/or the process conditions of forming the web.

The temperature of the blend is elevated within the extruder 14 by a conventional heating arrangement (not shown) to melt the polymer and pressure is applied to the polymer by the pressure-applying action of a turning screw (not shown), located within the extruder, to form the polymer into an extrudable composition. Preferably the polymer is heated to a temperature of at least about 175 °C if polypropylene is utilized as the fiber forming polymer. The polymer is then forwarded by the pressure applying action of the turning screw to a fiber forming arrangement 16 which may, for example, be a conventional meltblowing die arrangement. Meltblowing die arrangements are described in US-A-3,978,185 to Buntin et al and US-A-3,849,241 to Buntin et al. The elevated temperature of the polymer is maintained in the fiber forming arrangement 16 by a conventional heating arrangement (not shown). The fiber-forming arrangement generally extends a distance in the cross-machine direction which may be about equal to the width of the fibrous porous nonwoven web which is to be formed by the process. The fiber-forming arrangement 16 extrudes and attenuates the fibers 18 and directs them onto a moving forming screen 20. Upon impacting the forming screen 20, the fibers 18 may, depending upon known process conditions, adhere to each other to form the fibrous porous web 22. If not, a nip roller 24, in combination with the forming screen 20 can act to make the web 22 self supporting. If desired, the web 22 may be passed through a thermal point bonding arrangement 26 including rollers 28 and 30 to consolidate the web 22 even further. The combination of elevated temperature and elevated pressure conditions which effect extrusion of the polymer will vary over wide ranges. For example, at higher elevated temperatures, lower elevated pressures will result in satisfactory extrusion rates and, at higher elevated pressures of extrusion, lower elevated temperatures will effect satisfactory extrusion rates.

During or shortly after formation of the fibrous porous web 22, the high hydrophile-lipophile surface active agent is sprayed onto the surface of the fibers forming the web 22. In many instances the heat of the molten fibers 18 cooling after extrusion will be sufficient to effect drying of the high hydrophile-lipophile balance surface active agent. However, in some instances, the web 22 will have to be passed through a heating arrangement 32 which can include heating cans 34 and 36 to effect drying. The heating can drying temperature will vary with the surface active agent and polymer utilized. In any event the drying conditions are to be adjusted so that at least about 0.05, weight percent of the resultant web 22, of surface active agent will be on the surface of the web 22. For example, from about 0.05 percent, by weight, to about 3 percent, by weight of the web 22 of surface active agent will be on the surface of the web 22. More particularly, from about 0.10 percent, by weight, to about 1.0 percent, by weight of the web 22, of surface active agent will be on the surface of the web 22. For example, from about 0.1 percent, by weight, to about 0.4 percent, by weight, of the web 22, of surface active agent will be on the surface of the web 22. Even more particularly, from about 0.20 percent, by weight, to about 0.30 percent by weight of the web 22, of surface active agent will be on the surface of the web 22.

Determination of the weight percentage of the surface active agent on the surface of the web at this point in the process can be determined by: (1) weighing the initial sample of material; (2) quantitatively extracting the surface active agent from the surface of the web 22 using an appropriate solvent; (3) determining the amount of surface active agent in the extraction solvent by such means as ultraviolet spectroscopy, infra-red spectroscopy, gravimetric analysis etc. (This may require making up a series of concentration standards of the surface active agent in the extracting fluid to calibrate the analytical equipment/method/technique. Manufactures of surface active agent often will supply methods for determining surface active agent quantitatively and qualitatively.); and (4) dividing the amount of surface active agent by the initial web 22 sample weight and multiplying by 100.

Once the high hydrophile-lipophile balance surface active agent has been applied to the surface of high hydrohead of the web 22, the web 22 is passed through the gaps of two conventional corona discharge units 38. The two corona units are arranged so one treats one side of the web 22 and the other corona unit treats the other side of the web 22. One desirable corona discharge unit can be obtained from Enercon Ind. Corporation under trade designation Model SS 1223. The gaps of the corona discharge treatment apparatus may be maintained at about 0.065 inches. Standard metal rolls are used as the ground electrode. The base metal ground electrode roll may be buffered with 1 wrap of 0.0127mm (0.5 mil) polyester to substantially prevent arcing of the corona unit and pinholing in the high hydrohead fibrous porous web 22. Such buffering

reduces the effectiveness of the corona discharge unit by approximately 20% for each wrap of 0.0127mm (0.5 mil) film used. The line speed of the high hydrohead web material 22 and the voltage and amperage of the corona discharge unit 38 are adjusted so that the equivalent of at least about 8.6 Wmin/m² (0.8 watt min. per ft²) per side of corona discharge is applied to the web material 22. For example, the equivalent of from about 8.6 to about 161.5 Wmin/m² (about 0.8 to about 15 watt min per ft²) per side of the web material 22 of corona discharge may be applied to the web material 22. Accordingly, the equivalent of from about 10.7 to about 107.6 Wmin/m² (1 to about 10 watt min. per ft²) per side of the web material 22 of corona discharge may be applied to the web material 22. More particularly, the equivalent of from about 21.53 to about 86.11 Wmin/m² (about 2 to about 8 watt min. per ft²) per side of the web material 22 of corona discharge may be applied to the web material 22.

Once the corona discharge unit 38 has applied the appropriate amount of charge to the web material 22, the web material 22, may be wound up on a storage roll 40. The corona treated web material 22 may later be used in a wide variety of applications which require or desire utilization of a material having acceptable retentive water absorbency and retentive water acquisition rates. This method of treating a high hydrohead fibrous porous web material 22 has been found to increase the web's retentive acquisition rate (averaged normalized rate of water absorption in subsequent reabsorptions as compared to the initial absorption rate) and retentive absorbency (averaged normalized amount of water absorbed in subsequent reabsorptions as compared to the amount initially absorbed).

Another embodiment is schematically illustrated in Figure 2. In this situation the surface active agent may be applied in neat form or from solution by any of a number of conventional application methods. Exemplary of which is dip-and-squeeze. The dip-and-squeeze method is illustrated in Figure 2 with the dip-and-squeeze apparatus 42 including a dipping bath 44 and a pair of squeezing rollers 46 and 48. In this process at least about 0.05%, by weight, of the web material 22 of high hydrophile-lipophile balance surface active agent is applied to the web material 22. For example, from about 0.05% to about 3%, by weight of the web material, of high hydrophile-lipophile balance surface active agent may be applied to the web material 22. Even more particularly, from about 0.1% to about 1%, by weight of the web material 22, of high hydrophile-lipophile balance surface active agent may be applied to the material 22. More particularly, from about 0.1% to about 0.4%, by weight of the web material 22, of high hydrophile-lipophile balance surface active agent may be applied to the web material 22. Even more particularly, from about 0.2% to about 0.3%, by weight of the web material 22, of high hydrophile-lipophile surface active agent may be applied to the web material 22. The remainder of the process is the same as the process described with respect to Figure 1.

Of course, other conventional methods can be used for the production of the nonwoven web 22.

EXAMPLE

In order to demonstrate the improved retentive water absorbency and improved retentive water acquisition rate of corona discharge treated high hydrophile-lipophile balance web materials of the invention, samples of commercially available wet wipers available from the Kimberly-Clark Corporation under the trademark Kimtex were treated in accordance with the teachings of the present invention. The wiper material was an approximate 62.82 g/m² (2 oz. per yd²) meltblown polypropylene material which had already been treated with a sufficient amount of Aerosol OT, Na-di(2-ethylhexyl) sulphosuccinate, to have a surface concentration of Aerosol OT of about 0.30 weight percent or about 0.2 g/m² (0.006 oz. per yd²). This web material was subjected to corona discharge treatment in accordance with our invention. The amount of corona discharge applied to the sample was varied by varying the line speed of the web material as it moved through the gaps of each of the two corona discharge electrodes. Each of the two electrodes were three feet in length and had their gap set at 0.17 cm (0.065 inch) and the power supply was set at 1.25 kilowatts for each of the two electrodes. The ground roll of each electrode was buffered with one wrap of 0.0127 mm (0.5 mil) polyester to prevent arcing and pinholing. As has been previously stated this buffering reduces the effectiveness of the corona discharge by about 20 percent. Samples were made with the line speed (ls) of the web set at 7.62, 15.24, 30.48, 91.44, 121.92 and 182.88 m (25, 50, 100, 300, 400 & 600 ft) per min. The corresponding watt-min per m²(per ft²) per side of corona discharge values are 143.2, 71.0, 35.5, 11.8, 8.93 and 5.92 (13.3, 6.6, 3.3, 1.1, 0.83 and 0.55), respectively. For example, the corona charge placed on each side of the 121.92 (400 feet) per minute sample can be calculated as follows: 1250 watts per side times 0.80 efficiency divided by 0.9144 m (3 feet) electrode length divided by 121.92 m (400 feet) per minute equals 8.99 Wmin/m² (0.83 watt min per ft²) per side.

Testing of these materials and samples of non-corona treated material was conducted in accordance with Test A. The results of this testing is reported below in the Table, below.

Sample I. D.	Total Sample Weight (g)	Actual Sample Weight (g)	Water Absorbed (g) 1.2 sec	Water Absorbed (g) 2.4 sec	Water Rate (g/sec)	Normalized Rate (g/g sec)	Water Absorbed (g) 1 min	Water Absorbed (g) 2 min	Normalized Water Absorbed (g water/g wiper) 1 min	Normalized Rate (g/g sec)	Average Normalized Rate (g/g sec)	Decrease in Average Normalized Water Absorbed (g) 1 min	Decrease in Average Normalized Water Absorbed (g) 2 min	In Average Normalized Water Absorbed in Grams At 2 Min. (%)	
UI - 1 (a)	1.20	1.17	0.50	1.00	0.417	0.357	3.20	3.80	2.74	3.26	0.398	MA	2.84	3.40	NA
UI - 2 (a)	1.08	1.05	0.45	0.90	0.417	0.398	3.12	3.70	2.98	3.53					NA
UI - 3 (a)	1.13	1.10	0.40	0.80	0.458	0.418	3.20	3.87	2.92	3.53					
UI - 4 (a)	1.13	1.10	0.40	0.80	0.458	0.418	2.97	3.62	2.71	3.30					
UI - 1 (b)	1.20	1.17	0.20	0.70	0.292	0.250	2.20	2.70	1.69	2.31	0.245	38	1.93	2.37	30
UI - 2 (b)	1.08	1.05	0.30	0.70	0.292	0.279	2.25	2.70	2.15	2.50					
UI - 3 (b)	1.13	1.10	0.23	0.68	0.283	0.258	2.25	2.78	2.03	2.53					
UI - 4 (b)	1.11	1.08	0.00	0.00	0.000	0.000	1.80	2.20	1.67	2.04					
UI - 1 (c)	1.20	1.17	0.00	0.00	0.000	0.000	0.00	0.00	0.00	0.00	0.000	88	0.93	1.18	67
UI - 2 (c)	1.10	1.07	0.00	0.20	0.083	0.078	1.50	1.90	1.41	1.78					
UI - 3 (c)	1.15	1.12	0.00	0.00	0.000	0.000	1.15	1.46	1.03	1.31					67
UI - 4 (c)	1.13	1.10	0.00	0.30	0.125	0.114	1.40	1.77	1.28	1.61					74
UI - 1 (d)	1.20	1.17	0.00	0.00	0.000	0.000	0.00	0.00	0.00	0.00	0.000	100	0.71	0.87	75
UI - 2 (d)	1.08	1.05	0.00	0.00	0.000	0.000	0.22	0.95	0.69	0.91					
UI - 3 (d)	1.13	1.10	0.00	0.00	0.000	0.000	1.15	1.40	1.05	1.28					
UI - 4 (d)	1.11	1.08	0.00	0.00	0.000	0.000	1.18	1.40	1.10	1.30					
25 - 1 (a)	0.92	0.89	0.60	1.10	0.458	0.516	3.10	3.70	3.49	4.17	0.478	MA	3.51	4.16	MA
25 - 2 (a)	0.99	0.96	0.60	1.10	0.458	0.479	3.15	3.75	3.29	3.92					
25 - 3 (a)	0.98	0.95	0.80	1.10	0.417	0.440	3.55	4.15	3.75	4.38					
25 - 1 (b)	0.95	0.92	0.95	1.20	0.500	0.545	3.15	3.70	3.44	4.03	0.536	0	3.28	3.86	7
25 - 2 (b)	0.97	0.94	0.75	1.15	0.479	0.511	2.95	3.50	3.15	3.74					
25 - 3 (b)	0.92	0.89	0.65	1.17	0.468	0.550	2.88	3.38	3.25	3.81					
25 - 1 (c)	0.92	0.89	0.68	1.08	0.450	0.507	2.84	3.35	3.20	3.78	0.470	2	3.00	3.56	14
25 - 2 (c)	0.98	0.95	0.68	1.05	0.458	0.463	2.70	3.22	2.85	3.40					
25 - 3 (c)	0.98	0.95	0.53	1.00	0.417	0.440	2.78	3.32	2.94	3.51					
25 - 1 (d)	0.95	0.92	0.50	0.85	0.354	0.386	2.30	2.77	2.51	3.02	0.458	4	2.76	3.30	21
25 - 2 (d)	0.91	0.88	0.49	1.00	0.417	0.475	2.51	2.99	2.36	3.41					
25 - 3 (d)	0.97	0.94	0.65	1.15	0.479	0.511	2.72	3.24	2.90	3.46					
50 - 1 (a)	1.10	1.07	0.80	1.20	0.500	0.449	3.50	4.20	3.28	3.94	0.396	MA	3.31	3.96	MA
50 - 2 (a)	1.08	1.05	0.60	1.00	0.417	0.398	3.42	4.10	3.27	3.92					
50 - 3 (a)	1.07	1.04	0.60	0.80	0.333	0.321	3.50	4.18	3.38	4.03					
50 - 1 (b)	1.03	1.00	0.60	1.10	0.458	0.459	3.10	3.72	3.11	3.73	0.448	6	2.92	3.51	12
50 - 2 (b)	1.07	1.04	0.63	1.13	0.471	0.456	2.84	3.44	2.74	3.32					
50 - 3 (b)	1.10	1.07	0.60	1.10	0.458	0.429	3.10	3.70	2.91	3.47					
50 - 1 (c)	1.11	1.08	0.61	1.09	0.454	0.422	2.62	3.40	2.62	3.16	0.432	0	2.67	3.20	19
50 - 2 (c)	1.08	1.05	0.62	1.20	0.500	0.478	2.75	3.41	2.63	3.26					
50 - 3 (c)	1.08	1.05	0.63	1.00	0.417	0.398	2.88	3.33	2.75	3.18					
50 - 1 (d)	1.10	1.07	0.53	0.90	0.375	0.351	2.50	3.02	2.34	2.83	0.372	6	2.42	2.96	27
50 - 2 (d)	1.05	1.02	0.45	0.90	0.375	0.345	2.45	3.06	2.45	3.01					
50 - 3 (d)	1.03	1.00	0.47	0.95	0.396	0.397	2.50	3.06	2.51	3.05					

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Sample I. O.	Total Sample Weight (g)	Actual Sample Weight (g)	Water Absorbed (g) 1.2 sec	Water Absorbed (g) 2.4 sec	Rate (g/sec)	Normalized Rate (g/g sec)	Water Absorbed (g) 1 min	Water Absorbed (g) 2 min	Normalized Water Absorbed (g water/g wiper) 1 min	Normalized Water Absorbed (g water/g wiper) 2 min	Average Normalized Rate (g/g sec)	Decrease in Average Normalized Rate (%)	Average Normalized Water Absorbed (g) 1 min	Average Normalized Water Absorbed (g) 2 min	Decrease in Average Normalized Water Absorbed in Grams At 1 Min. (%)	Decrease in Average Normalized Water Absorbed in Grams At 2 Min. (%)
100 - 1 (a)	1.02	0.99	0.65	1.08	0.450	0.456	3.42	4.08	3.47	4.13	0.469	MA	3.42	4.07	MA	MA
100 - 2 (a)	1.10	1.07	0.65	1.05	0.450	0.450	3.52	4.20	3.30	3.94						
100 - 3 (a)	1.07	1.04	0.72	1.32	0.550	0.530	3.63	4.30	3.50	4.15						
100 - 1 (b)	1.05	1.02	0.75	1.20	0.500	0.492	3.10	3.65	3.05	3.59	0.428	9	2.88	3.41	16	16
100 - 2 (b)	1.09	1.06	0.51	0.81	0.338	0.320	2.70	3.23	2.55	3.06						
100 - 3 (b)	1.00	0.97	0.30	1.10	0.658	0.474	2.93	3.48	3.03	3.60						
100 - 1 (c)	1.02	0.99	0.40	0.90	0.375	0.380	2.38	3.40	2.92	3.44						
100 - 2 (c)	1.10	1.07	0.40	0.92	0.383	0.359	2.82	3.35	2.64	3.14						
100 - 3 (c)	1.05	1.02	0.55	1.10	0.458	0.450	2.77	3.32	2.72	3.26						
100 - 1 (d)	1.00	0.97	0.40	0.90	0.375	0.388	2.60	3.10	2.69	3.21	0.411	12	2.65	3.15	23	23
100 - 2 (d)	1.08	1.05	0.40	0.92	0.383	0.364	2.65	3.15	2.53	3.01						
100 - 3 (d)	1.02	0.99	0.57	1.15	0.479	0.485	2.68	3.20	2.72	3.24						
300 - 1 (a)	1.01	0.98	0.80	1.31	0.546	0.559	3.30	3.90	3.38	3.99	0.476	MA	3.30	3.96	MA	MA
300 - 2 (a)	1.00	0.97	0.53	1.08	0.450	0.465	3.15	3.75	3.26	3.88						
300 - 3 (a)	0.98	0.95	0.52	0.92	0.383	0.404	3.10	3.80	3.27	4.01						
300 - 1 (b)	0.96	0.93	0.64	1.10	0.658	0.494	2.52	3.00	2.72	3.24	0.547	0	2.95	3.47	11	12
300 - 2 (b)	0.99	0.96	0.80	1.40	0.583	0.609	3.01	3.52	3.15	3.68						
300 - 3 (b)	1.00	0.97	0.80	1.25	0.521	0.539	2.88	3.38	2.98	3.50						
300 - 1 (c)	1.00	0.97	0.00	0.68	0.258	0.267	2.28	2.75	2.36	2.84						
300 - 2 (c)	1.00	0.97	0.00	0.80	0.333	0.344	2.45	2.97	2.53	3.07						
300 - 3 (c)	0.98	0.95	0.30	1.00	0.417	0.440	2.47	2.94	2.61	3.10						
300 - 1 (d)	1.00	0.97	0.40	0.90	0.375	0.388	2.48	2.99	2.56	3.09	0.449	6	2.72	3.23	18	18
300 - 2 (d)	0.98	0.95	0.30	1.02	0.425	0.449	2.66	3.12	2.81	3.29						
300 - 3 (d)	0.95	0.92	0.54	1.12	0.467	0.509	2.56	2.94	2.79	3.32						
400 - 1 (a)	1.08	1.05	0.72	1.25	0.521	0.498	3.35	3.95	3.20	3.77	0.480	MA	3.23	3.83	MA	MA
400 - 2 (a)	1.08	1.05	0.70	1.32	0.550	0.525	3.42	4.04	3.27	3.86						
400 - 3 (a)	1.08	1.05	0.50	1.05	0.438	0.418	3.37	4.04	3.22	3.86						
400 - 1 (b)	1.08	1.05	0.23	0.80	0.333	0.318	2.50	3.10	2.39	2.96	0.276	43	2.30	2.82	29	26
400 - 2 (b)	1.06	1.03	0.00	0.55	0.229	0.223	2.15	2.67	2.09	2.60						
400 - 3 (b)	1.08	1.05	0.14	0.72	0.360	0.287	2.52	3.02	2.41	2.88						
400 - 1 (c)	1.08	1.05	0.52	0.94	0.392	0.374	2.24	2.82	2.14	2.50	0.209	56	1.77	2.18	45	45
400 - 2 (c)	1.08	1.05	0.00	0.00	0.000	0.000	1.22	1.62	1.17	1.55						
400 - 3 (c)	1.10	1.07	0.00	0.65	0.271	0.254	2.15	2.65	1.91	2.48						
400 - 1 (d)	1.05	1.02	0.45	0.80	0.353	0.327	2.50	2.97	2.46	2.92	0.306	36	2.21	2.66	32	31
400 - 2 (d)	1.05	1.02	0.40	0.95	0.396	0.369	2.31	2.77	2.27	2.72						
400 - 3 (d)	1.09	1.06	0.80	0.51	0.212	0.201	2.01	2.46	1.90	2.35						

Sample	Total Sample Weight (g)	Actual Sample Weight (g)	Water Absorbed (g) 1.2 sec	Water Absorbed (g) 2.4 sec	Rate (g/sec)	Normalized Rate (g/g sec)	Water Absorbed (g) 1 min	Water Absorbed (g) 2 min	Normalized Water Absorbed (g water/g wiper) 1 min	Normalized Water Absorbed (g water/g wiper) 2 min	Average Normalized Rate (g/g sec)	Decrease in Average Normalized Rate (%)	Averaged Water Absorbed (g) 1 min	Averaged Water Absorbed (g) 2 min	Decrease in Average Normalized Water Absorbed in Grams At 1 Min. (%)	Increase in Average Normalized Water Absorbed in Grams At 2 Min. (%)
600 - 1 (a)	1.05	1.02	0.75	1.20	0.500	0.492	3.12	3.74	3.07	3.68	0.459	NA	3.22	3.81	NA	NA
600 - 2 (a)	1.05	1.02	0.55	1.05	0.458	0.431	3.35	3.97	3.29	3.90						
600 - 3 (a)	1.02	0.99	0.68	1.08	0.450	0.456	3.26	3.60	3.30	3.85	0.595	14	2.65	3.18	18	17
600 - 1 (b)	1.01	0.98	0.39	0.99	0.412	0.422	2.80	3.31	2.87	3.39						
600 - 2 (b)	1.01	0.98	0.29	0.89	0.371	0.380	2.58	3.10	2.64	3.17						
600 - 3 (b)	1.01	0.98	0.40	0.90	0.375	0.386	2.40	2.91	2.46	2.98						
600 - 1 (c)	1.05	1.02	0.00	0.00	0.000	0.000	0.00	0.00	0.00	0.00	0.029	94	1.02	1.31	68	66
600 - 2 (c)	1.00	0.97	0.00	0.05	0.021	0.022	1.45	1.87	1.50	1.93						
600 - 3 (c)	1.00	0.97	0.00	1.15	0.062	0.064	1.52	1.92	1.57	1.99						
600 - 1 (d)	1.00	0.97	0.00	0.00	0.000	0.000	0.00	0.21	0.00	0.22	0.053	93	1.01	1.37	69	64
600 - 2 (d)	1.01	0.98	0.00	0.00	0.000	0.000	0.95	1.30	0.97	1.33						
600 - 3 (d)	1.00	0.97	0.00	0.45	0.095	0.098	2.00	2.40	2.07	2.56						

The data of Table I may be interpreted as follows: (1) UT represents the non-corona treated samples whereas the 25, 50, 100, 300, 400, 600 represents the feet (1 foot = 0.3045 m) per minute of the web as it passed through the corona discharge gap; (2) three replicate samples of each treated material were taken with each of these being represented by the number 1 or 2 or 3; (3) each of the samples was subjected to testing in accordance with Test A four times with the first test being designated by the letter (a) the second

represented by the letter (b) the third being represented by the letter (c) and the fourth being represented by the letter (d). Thus 600-3(d) stands for the results of the fourth time the third replicate sample of material treated at 182.7 m (600 feet) per minute was tested in accordance with Test A.

From the above results reported in Table I, it is clear that materials treated in accordance with our invention line speeds of about 121.92 m (400 feet) per minute or less (about 8.6 Wmin/m² (0.8 watt-min per ft²) per side or greater) have significantly smaller decreases in average normalized rate of water acquisition and in averaged normalized water absorbed at one minute and two minutes. Such materials can be repeatedly reused as wiper materials helping both the environment because of less wipers being used and the user because less materials may be purchased.

Claims

1. A method of preparing a high hydrohead fibrous porous web with improved retentive absorption comprising the steps of:
 - providing a high hydrohead fibrous porous web having a surface concentration of at least about 0.05 percent, by weight of the web, of a surface active agent having a hydrophile-lipophile balance of at least about 6; and
 - applying a corona discharge equivalent to a charge of at least about 8.6 Wmin/m² (about 0.8 watt minute per square foot) per side of the web to the surface active agent bearing web;
 - wherein the resulting web has a percent decrease in the averaged normalized water absorbed, at two minutes, of less than about 50, preferably 25 weight percent in each of the second, third and fourth times the material is tested in accordance with absorbency test A as explained in the description, when compared to the averaged normalized water absorbed upon being initially tested in accordance with absorbency test A.
2. The method of claim 1, wherein the resulting web has a percent decrease in the averaged normalized rate of water absorbed in the first 2.4 seconds of absorption, of less than about 50 percent, preferably less than about 25 percent, more preferably less than about 10 percent in each of the second, third and fourth times the material is tested in accordance with absorbency test A when compared to the averaged normalized rate of water absorbed upon being initially tested in accordance with absorbency test A.
3. The method of claim 1 or 2, wherein the surface concentration of the surface active agent is from about 0.05% to about 3%, preferably from about 0.1% to about 1%, more preferably from about 0.1% to about 0.4% percent, particularly from about 0.2% to about 0.3%, by weight of the web material.
4. The method of one of the preceding claims wherein the equivalent of from about 8.6 to about 161.3 Wmin/m² (about 0.8 to about 15 watt minute per square foot), preferably from about 10.7 to about 107.5 Wmin/m² (about 1 to about 10 watt minute per square foot), more preferably from about 21.5 to about 86.0 Wmin/m² (about 2 to about 8 watt minute per square foot) per side of the web material of corona discharge is applied to the web material.
5. The method of one of the preceding claims, wherein the surface active agent is selected from the group consisting of one or more wetting agents, emulsions, emulsifiers, detergents and dispersants.
6. The product preparable by the method of one of claims 1 to 5.

Patentansprüche

1. Verfahren zur Herstellung einer porösen Faserbahn hoher "Hydrohead"-Zahl mit verbesserter bleibender Absorptionfähigkeit, die folgenden Schritte umfassend:
 - Schaffung einer porösen Faserbahn hoher "Hydrohead"-Zahl mit einer nach dem Gewicht der Bahn bemessenen Oberflächenkonzentration von mindestens etwa 0,05 Prozent eines oberflächenaktiven Mittels mit einem Hydrophile-Lipophile-Gleichgewicht von mindestens etwa 6; und
 - Anwenden einer Corona-Entladung, entsprechend einer Ladung von mindestens 8,6 Wmin/m² (etwa 0,8 Wattminuten pro Quadratfuß) pro Bahnseite auf die Bahn mit dem oberflächenaktiven Mittel;
 - wobei die resultierende Bahn jeweils nach zwei Minuten beim zweiten, dritten und vierten Mal, zu dem das Material gemäß dem Absorptionstest A wie in der Beschreibung ausgeführt getestet wird, einen

anteilmäßigen Rückgang des durchschnittlichen normalisierten absorbierten Wassers von weniger als 50, vorzugsweise 25 Gewichtsprozent, aufweist im Vergleich zu dem durchschnittlichen normalisierten Wasser, das nach einem ersten Test gemäß dem Absorptionstest A absorbiert wurde.

- 5 2. Verfahren nach Anspruch 1, wobei die resultierende Bahn in den ersten 2,4 Sekunden der Absorption jeweils beim zweiten, dritten und vierten Mal, zu dem das Material gemäß dem Absorptionstest A getestet wird, einen anteilmäßigen Rückgang der durchschnittlichen normalisierten Wasserabsorptionsgeschwindigkeit von weniger als 50 Prozent, vorzugsweise weniger als etwa 25 Prozent, insbesondere von weniger als etwa 10 Prozent, aufweist im Vergleich zu der durchschnittlichen normalisierten Wasserabsorptionsgeschwindigkeit nach einem ersten Test gemäß dem Absorptionstest A.
- 10 3. Verfahren nach Anspruch 1 oder 2, wobei die Oberflächenkonzentration des oberflächenaktiven Mittels, gemessen am Gewicht des Bahnmaterials, zwischen etwa 0,05% und etwa 3% liegt, vorzugsweise zwischen etwa 0,1% und etwa 1%, insbesondere zwischen etwa 0,1% und etwa 0,4% und wiederum insbesondere zwischen etwa 0,2% und etwa 0,3%.
- 15 4. Verfahren nach einem der oben genannten Ansprüche, wobei pro Seite des Bahnmaterials eine Corona-Entladung entsprechend zwischen etwa 8,6 und etwa 161,3 Wmin/m² (etwa 0,8 bis etwa 15 Wattminuten pro Quadratfuß), vorzugsweise zwischen etwa 10,7 und etwa 107,5 Wmin/m² (etwa 1 bis etwa 10 Wattminuten pro Quadratfuß), insbesondere zwischen etwa 21,5 und etwa 86,0 Wmin/m² (etwa 2 bis etwa 8 Wattminuten pro Quadratfuß) auf dem Bahnmaterial angewendet wird.
- 20 5. Verfahren nach einem der vorhergehenden Ansprüche, wobei das oberflächenaktive Mittel ausgewählt ist aus der Gruppe bestehend aus einem oder mehreren Benetzungsmitteln, Emulsionen, Emulgatoren, Detergentien und Dispergiernmitteln.
- 25 6. Produkt, welches durch das Verfahren eines der Ansprüche 1 bis 5 herstellbar ist.

Revendications

- 30 1. Procédé de préparation d'une nappe poreuse fibreuse à fort "indice de colonne d'eau" ayant une capacité d'absorption rétentive améliorée, ledit procédé comprenant :
 - la fourniture d'une nappe poreuse, fibreuse, à fort "indice de colonne d'eau" ayant une concentration superficielle d'au moins environ 0,05% par rapport au poids de la nappe, d'un agent surfactif ayant une balance hydrophile-lipophile d'au moins environ 6 ;
 - 35 - l'application d'une décharge de couronne équivalente à une charge d'au moins environ 8,6 Wmin/m² (environ 0,8 watt minute par pied²), par face de la nappe, à la nappe portant ledit agent surfactif,
 procédé dans lequel la nappe résultante offre une réduction de la quantité moyenne et normalisée d'eau absorbée, en deux minutes, inférieur à environ 50%, de préférence de 25%, en poids au cours de chacune des deuxième, troisième et quatrième fois que le matériau est soumis au test de capacité d'absorption A tel qu'explicité dans la description, par comparaison avec la quantité moyenne et normalisée d'eau absorbée lorsque la nappe est soumise pour la première fois audit test de capacité d'absorption A.
- 40 2. Procédé selon la revendication 1, dans lequel la nappe résultante offre une réduction de la quantité moyenne et normalisée d'eau absorbée, au cours des premières 2,4 secondes d'absorption, de moins d'environ 50%, de préférence de moins d'environ 25%, et mieux de moins d'environ 10% en poids au cours de chacune des deuxième, troisième et quatrième fois que le matériau est soumis au test de capacité d'absorption A, par comparaison avec la quantité moyenne et normalisée d'eau absorbée lorsque la nappe est soumise pour la première fois audit test de capacité d'absorption A.
- 50 3. Procédé selon la revendication 1 ou 2, dans lequel la concentration superficielle dudit agent surfactif est comprise entre environ 0,05% et environ 3%, de préférence entre environ 0,1% et environ 1%, mieux entre environ 0,1% et environ 0,4%, et mieux encore d'environ 0,2% à environ 0,3%, en poids du matériau de la nappe.
- 55

4. Procédé selon l'une des revendications précédentes, dans lequel il est appliqué au matériau de la nappe une décharge de couronne équivalente à une charge d'environ 8,6 à environ 161,3 Wmin/m² (environ 0,8 à environ 15 watts minute par pied²), de préférence d'environ 10,7 à environ 107,5 Wmin/m² (environ 1 à environ 10 watts minute par pied²), mieux d'environ 21,5 à environ 86,0 Wmin/m² (environ 2 à environ 8 watts minute par pied²) par face du matériau en nappe.
5. Procédé selon l'une des revendications précédentes, dans lequel ledit agent surfactif est choisi dans le groupe comprenant un ou plusieurs agents mouillants, émulsions, émulsifiants, détergents et dispersants.
6. Produit pouvant être préparé par le procédé selon l'une quelconque des revendications 1 à 5.

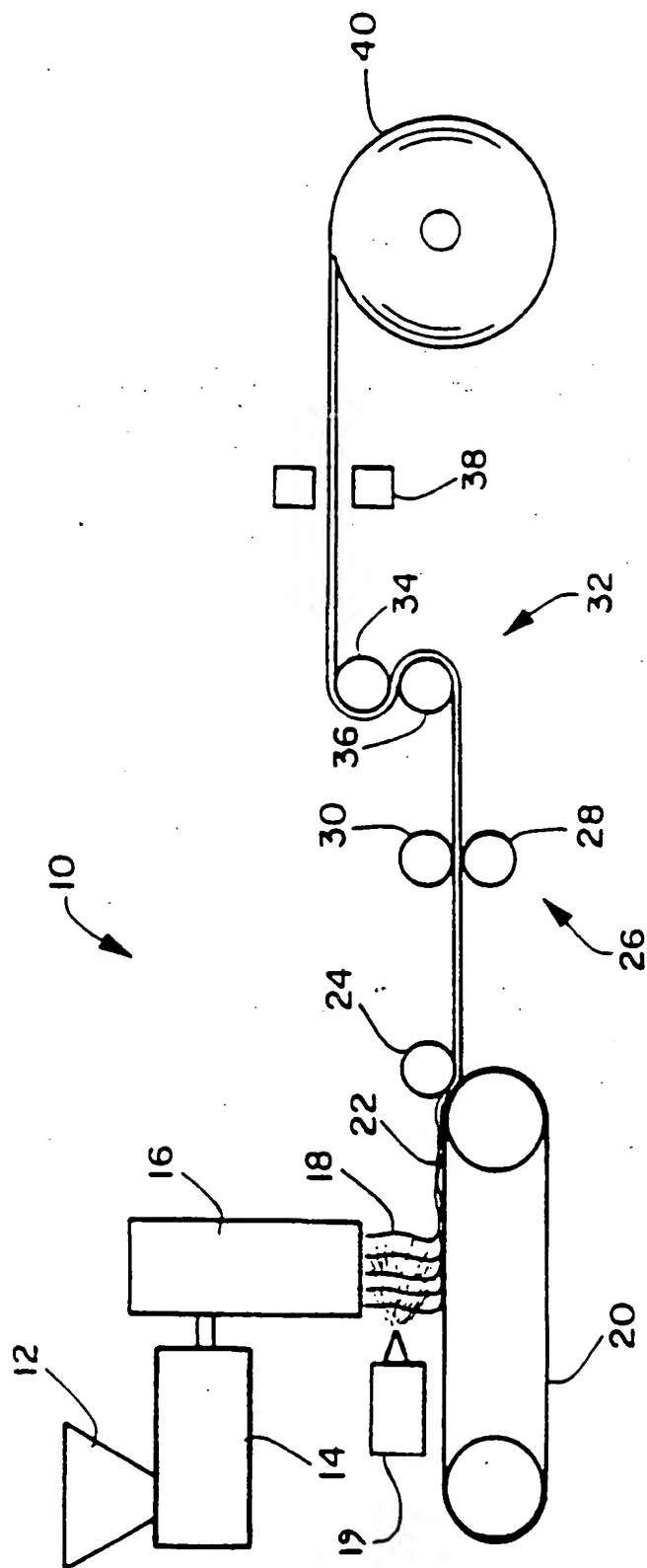


FIG. 1

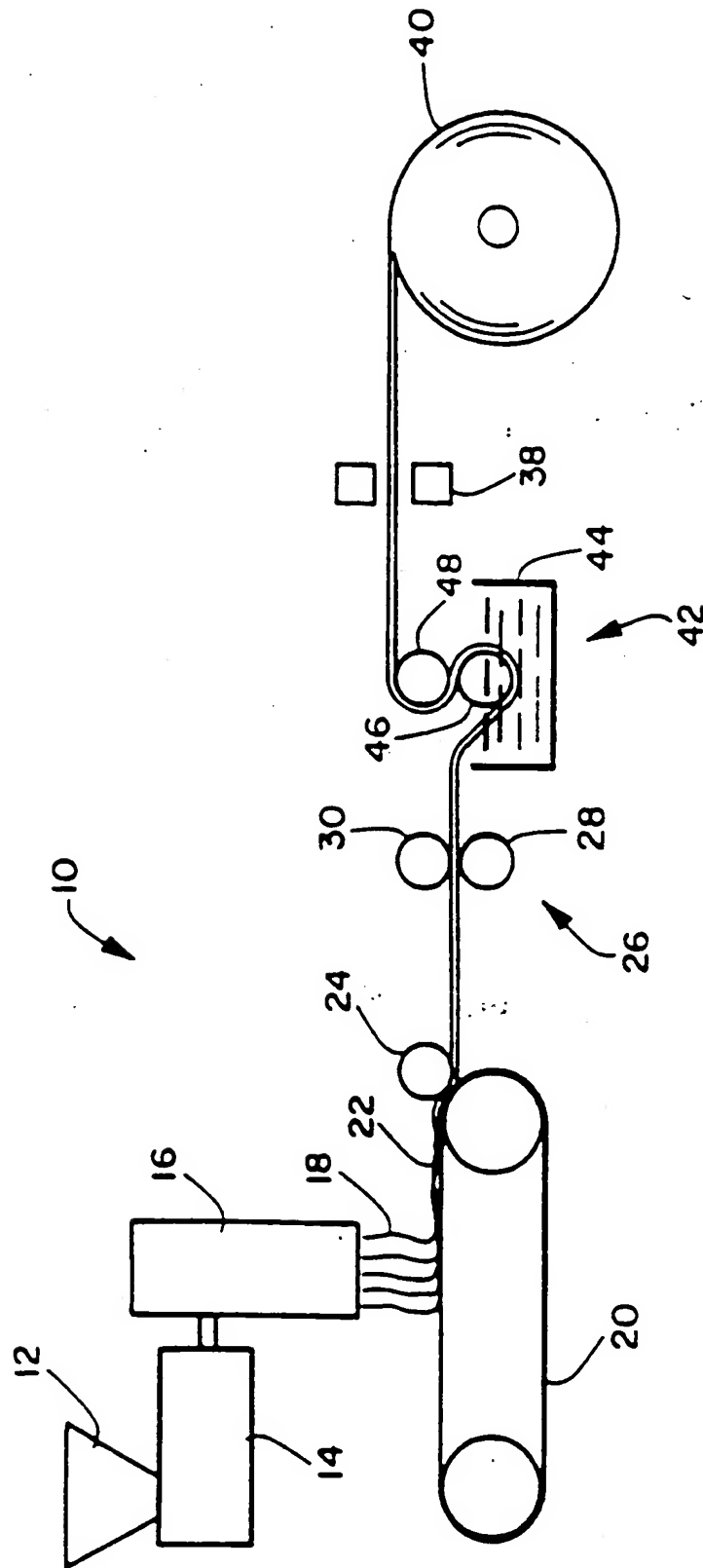


FIG. 2

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